

Case report: Ewing's sarcoma (ES) is the 2nd primary bone tumor of childhood, mostly located in the lower extremities. The incidence of primary cranial ES is <1%. Our patient is 9 years old female who has intracranial primary ES extending from the sphenoid bone corpus to the clivus border. This is a rare case of childhood that's originating from the sphenoid bone and spreading to such a very large intracranial area. Our aim is to provide data on the clinical and therapeutic course of a rare case.

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SUPPORTIVE CARE AND PALLIATIVE CARE

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ACTINOMYCES ODONTOLYTICUS: A RARE CAUSE OF PEDIATRIC FEBRILE NEUTROPENIA

Meric KAYMAK CIHAN¹,
Sonay İNCESÖY ÖZDEMİR¹, Belgin GÜLHAN²,
Arzu YAZAL ERDEM¹, Derya ÖZYÖRÜK¹,
Neriman SARI¹, İnci ERGÜRHAN İLHAN¹

¹ ANKARA CITY HOSPITAL, DEPARTMENT OF PEDIATRIC ONCOLOGY

² ANKARA CITY HOSPITAL, DEPARTMENT OF PEDIATRIC INFECTION DISEASES

Case report: Actinomyces spp. are gram-positive bacilli found in humans as a common flora of the oropharynx, gastrointestinal tract, and urogenital tract. We describe a case of Actinomyces odontolyticus bacteremia in an Ewing sarcoma and febrile neutropenic girl. This is the first time that bacteremia due to A. odontolyticus has been reported in a pediatric cancer patient. This case suggests that A. odontolyticus should be regarded as a possible cause of bacteremia in neutropenic pediatric cancer patients.

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TUMOR BIOLOGY, IMMUNOLOGY AND IMMUNOTHERAPY

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WITHDRAWN: THE EFFECT OF NIVOLUMAB IN PEDIATRIC MALIGNANT TUMORS: A SINGLE CENTER EXPERIENCE WITH EIGHT PATIENTS

Veysel GÖK¹, Firdevs AYDIN¹, Alper ÖZCAN¹,
Ebru YILMAZ¹, Ekrem UNAL¹,
Musa KARAKUKCU¹, Türkan PATIROĞLU¹,

Mehmet Akif ÖZDEMİR¹, Filiz KARAMAN²,
Orhan GÖRÜKMEZ³, Özlem GÖRÜKMEZ³,
Atıl BİSGİN⁴

¹ Division of Pediatric Hematology and Oncology, Department of Pediatrics, Erciyes University

² Division of Pediatric Radiology, Department of Radiology, Erciyes University

³ Department of Medical Genetics, Bursa Yüksek İhtisas Training and Research Hospital

⁴ Department of Medical Genetics, Medical Faculty, Çukurova University

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A CASE DIAGNOSED WITH FOUR DIFFERENT TUMORS

Fatma Tuba YILDIRIM, Derya ÖZYÖRÜK,
Arzu YAZAL ERDEM, Selma ÇAKMAKCI,
Neriman SARI, Sonay İNCESÖY, İnci İLHAN

Ankara City Hospital

Case report: Chromosomal breakage syndromes are characterized by cancer predisposition. Here we present a 27-month-old female with Fanconi Aplastic Anemia diagnosed with 4 tumors. Imaging showed brain mass causing the shift, liver mass and left kidney mass. She had diagnosed with high grade intracranial tm, wilms tm and hepatocellular ca. Because of refractory pancytopeni, she underwent HSCT. After 2months she developed intracranial embryonal tumor. The patient died with progression. Genetic tests revealed no mutation.

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